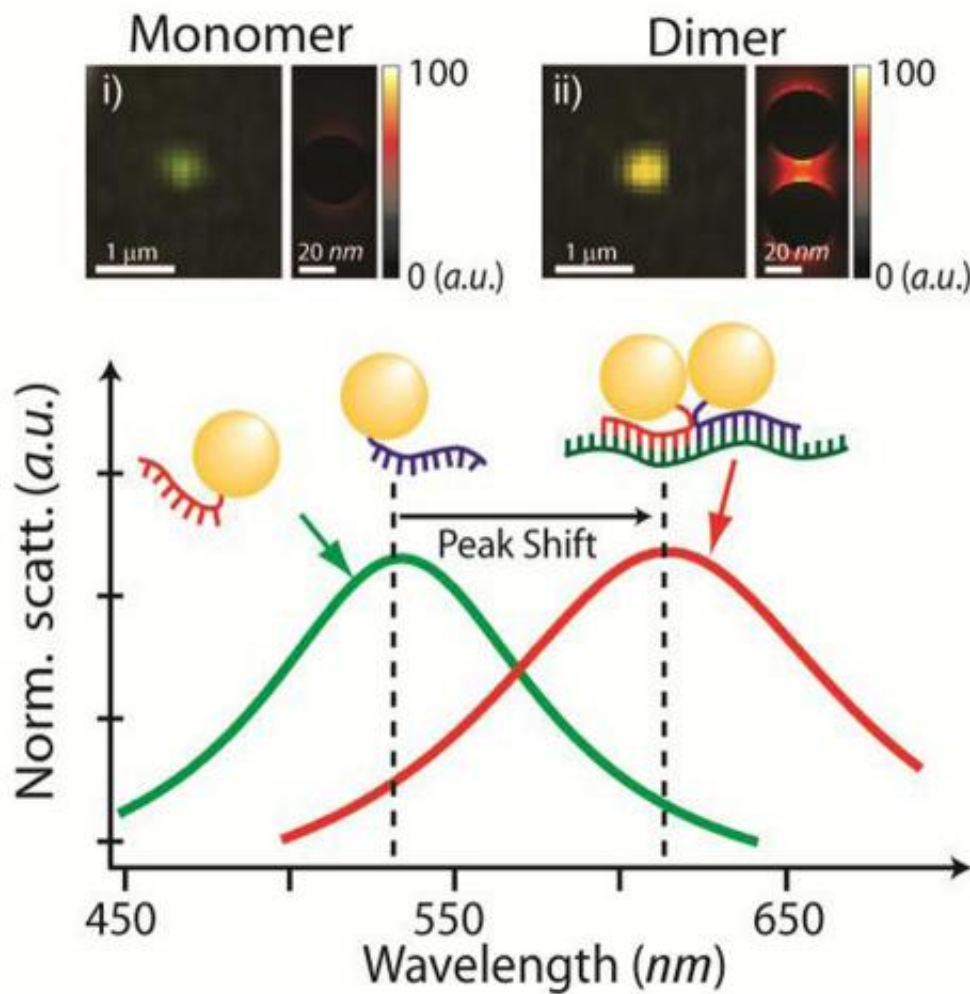


Gold nanoparticles help target, quantify breast cancer gene segments in a living cell

April 23 2014, by Natalie Van Hoose



A single gold nanoparticle, or monomer, appears green when illuminated (top left), while a pair of gold nanoparticles bound to an mRNA splice variant, or dimer, appears reddish (top right). Monomers and dimers also scatter light differently, as shown in the graph above. Credit: Purdue University image / Joseph Irudayaraj

(Phys.org) —Purdue University researchers have developed a way to detect and measure cancer levels in a living cell by using tiny gold particles with tails of synthetic DNA.

A team led by Joseph Irudayaraj, professor of agricultural and biological engineering, used gold nanoparticles to target and bind to fragments of [genetic material](#) known as BRCA1 messenger RNA splice variants, which can indicate the presence and stage of breast cancer. The number of these mRNA splice variants in a cell can be determined by examining the specific signal that light produces when it interacts with the gold nanoparticles.

"This is a simple yet sophisticated technique that can be used to detect cancer in a single cell and determine how aggressive it is," said Irudayaraj, who is also the deputy director of the Bindley Bioscience Center. "Being able to quantify these genetic molecules could ultimately help clinicians provide better and more individualized treatment to cancer patients."

The technique also could increase our understanding of cell biology and paves the way for genetic profiling and diagnosis based on a single cell, Irudayaraj said.

BRCA1 is a [tumor suppressor gene](#) that can transform a cell into a cancerous type under certain circumstances. Measuring the number of BRCA1 mRNA splice variants in a cell can indicate if the gene is being under-expressed, a possible sign of [breast cancer](#).

But current methods of detecting cancer rely on samples made up of hundreds or thousands of [cells](#) and cannot provide detailed information about how genes tied to cancer are being expressed in individual cells.

Irudayaraj and his team are the first to detect and quantify BRCA1 mRNA splice variants - fragments of genetic material that are removed when mRNA is formed - in a single cell. Splice variants can determine the fate of a cell and how specific proteins are expressed. Errors in the splicing process have been linked to a variety of diseases.

"With this method, we are basically able to spot a needle in a haystack - and we can determine if there are five needles in that haystack or if there are 50," he said.

Irudayaraj and his then-graduate research assistant, Kyuwan Lee, who is the first author of the study, adapted common nanotechnology methods to tackle the challenge of pinpointing mRNA splice variants in a living cell. They fabricated [gold nanoparticles](#) - more than 1,000 times smaller than the diameter of a human hair - and tagged them with strands of DNA complementary to BRCA1 mRNA splice variants.

When injected into a cell, the nanoparticles attached to either end of mRNA splice variants, forming structures known as dimers - each "like a couple holding hands," Irudayaraj said.

Because dimers give off a unique signal in the presence of light, the researchers could measure the number of dimers by illuminating the cell with a simple light source. The number of dimers corresponded to the number of BRCA1 mRNA splice variants in a cell.

Light behaves differently when it shines on a single gold particle, allowing the researchers to differentiate between dimers and free-floating [gold particles](#).

The researchers used two methods to quantify the dimers: spectroscopy, which measures the way light scatters when it encounters an object, and a colorimetric image on which [dimers](#) show as reddish dots while single

gold particles appear green.

The technique can quantify mRNA splice variants in a single cell in about 30 minutes.

Irudayaraj is modifying the system to speed up the process so that it can be used in tissue biopsies.

"If we can quantify key mRNA at single cell resolution in a tissue biopsy, that will be very powerful in terms of refining treatment protocols for key diseases," he said.

More information: "Quantitative imaging of single mRNA splice variants in living cells." Kyuwan Lee, Yi Cui, Luke P. Lee & Joseph Irudayaraj. *Nature Nanotechnology* (2014) [DOI: 10.1038/nnano.2014.73](https://doi.org/10.1038/nnano.2014.73). Received 22 November 2012 Accepted 11 March 2014 Published online 20 April 2014

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